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(50 Title: DEVICE FOR SITE DIRECTED NE	OVASC	ULARIZATION AND METHOD FOR SAME

(54) Title: DEVICE FOR SITE DIRECTED NEOVASCULARIZATION AND METHOD FOR SAME

(57) Abstract

The invention includes a device and method. The device is a site directed neovascularization device. The device in the site of the support. The device also includes a biological response modifier for inducing neovascularization. The biological responses modifier is adsorbed to the biocompatible support. The method is for directing in vivo neovascularization. The method requires adsorbing a biological response modifier for inducing neovascularization onto a biocompatible support. The step of contacting a therapertically effective amount of the adsorbed biological response modifier of at least one selected tissue then occurs. The method then involves directing neovascular cell growth at the contacted, selected tissue for a sufficient time to obtain a vascular structure. The method of this invention is useful for developing artificial organs and other tissues including nerves in an organism, and for sampling of cells and re-implantation after genetically altering the cells to produce a desired product.

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1	DEVICE FOR SITE DIRECTED NEOVASCULARIZATION
2	AND METHOD FOR SAME
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3	BACKGROUND OF THE INVENTION
4	1. Field of the Invention
5	The invention relates to a device and method for
6	directing the formation of new blood vessels and
7	artificial organs. Specifically, the invention relates
8	to a device and method for directing neovascularization
9	with a biological response modifier adsorbed onto a
10	support.
11	2. Description of the Background Art
	by Description of the Buonground Art
12	Angiogenesis is the formation of blood vessels in
13	<u>situ</u> and involves the orderly migration, proliferation,
14	and differentiation of vascular cells and occurs during
15	development. Angiogenesis is an infrequent event in the

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adult and is associated in adults with wound and fracture 1 2 repair. Exceptions to this are found in the female 3 reproductive system where this process occurs in the 4 follicle during development, in the corpus luteum during ovulation, and in the placenta during pregnancy. These 5 6 specific periods of angiogenesis are relatively brief and 7 highly regulated in contrast to the angiogenic events 8 associated with tumor growth and diabetic retinopathy. g The endothelial cell is considered to be the primary 10 cellular target for angiogenesis. Research efforts have 11 concentrated on the identity of polypeptide factors that 12 control endothelial cell proliferation. 13 heparin-binding growth factor (HBGF) 14 polypeptides has gained general acceptance as initiators 15 of angiogenesis especially during development.

The gene family for producing the heparin-binding growth factor family of polypeptides includes HBGF-1 (acidic fibroblast growth factor). HBGF-2 (basic fibroblast growth factor), and three additional HBGF-like structures, hst/KS, int-2, and FGF-5, each of which is encoded by an oncogene. The prototype HBGF polypeptides are potent inducers of endothelial cell migration and/or proliferation in vitro and are known to modulate the expression of endothelial cell derived proteases. Further, HBGF-1 and HBGF-2 are tightly adsorbed to the extracellular matrix presumably by their avid affinity for the glycosaminoglycan heparin. The between the HBGF prototypes and heparin protect these polypeptides from proteolytic modification. It has been suggested that the extracellular matrix can be the major source of HBGF-1 and HBGF-2 and activation can require hydrolytic extraction from sites of attachment for biological activity.

1 Hayek, et al (1987) reported the in vivo effect of 2 fibroblast growth factor in rat kidney. (Biochem. 3 Biophys. Res. Commun. 147:876-880.) The initiation of 4 angiogenesis by the direct stimulation of endothelial 5 cell proliferation is presumed to be a result of the 6 Class I heparin-binding growth factor (HBGF-I) and the Class II heparin-binding growth factor (HBGF-II). 7 8 polypeptides are potent endothelial cell growth factors 9 in vitro and angiogenesis signals in vivo. 10 polypeptides exert their biological response in vivo 11 through high affinity cell surface receptors. The HBGF-I 12 and HBGF-II share a structural similarity of 55 percent 13 and both are synthesized as polypeptides lacking an 14 apparent signal peptide sequence. Human cells which 15 express the HBGF-I mRNA transcript do not secrete the 16 polypeptide in vitro. Further, HBGF-II has been shown to 17 be associated with the extracellular matrix and heparin protects HBGF-I from proteolytic modification by plasmin. 18

19 PCT International Publication Number WO 87/01728 20 discloses recombinant fibroblast growth factors. 21 growth factors are examples of biological response 22 modifiers. This disclosure identifies the importance of 23 the growth factors for constructing vascular systems in 24 healing tissues. The invention of this disclosure is 25 directed to recombinant DNA sequences for encoding bovine 26 and human acidic and basic FGF and vectors bearing these 27 DNA sequences. This publication does not disclose a 28 device or method for site directed neovascularization.

The article, Van Brunt, et al., "Growth Factors

Speed Wound Healing", <u>Biotechnology</u> 6 (1988):25-30,

discloses the usefulness of growth factors in the

angiogenesis of damaged tissue. This article discloses a 1 2 sponge implant model for wound healing in animals. sponge consists of an inert polyvinyl alcohol that is 3 4 implanted under the skin of the animal. Growth factor is 5 then injected directly into the sponge. undergoes rapid healing and an increase in blood vessels 6 7 occurs at the wound site. The blood vessels resulting 8 from this invention do not form complete, permanent 9 vascular structures that are directed by a support to 10 which the growth factor is adsorbed. This article does not disclose a device or method for site directed 11 12 neovascularization.

13 U.S. Patent Number 4,699,141 to Lamberton, et al. discloses a container and method for neovascularization. 14 15 This invention has a sponge body that is throughout with a solution of fibrinogen and heparin. 16 17 The sponge body is placed adjacent to or around a 18 noncapillary blood vessel. Capillaries then grow into 19 the sponge. The sponge can then be used as a receptacle 20 for desired cells such as pancreas cells. This patent 21 does not disclose a device or method wherein the growth 22 of blood vessels is directed in a specific direction or 23 between specific sites. Neither the heparin nor collagen 24 in this invention modify a biological response. Both the 25 heparin and collagen are substrates upon 26 biological response modifier acts. The capillary growth 27 developed by this invention is a result 28 inflammatory response of the vessel to a foreign body or 29 the sponge. The blood vessels of this invention are not directed in their growth and do not form permanent 30 31 structures or long term structures. These blood vessels

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1 are not permanent because the fibrinogen support is absorbed by the organism before maturation of the blood 2 3 vessels can occur.

The blood vessels developed by the Lamberton, et al. invention are, essentially, a bundle of cells or This invention is capillaries within a sponge. identified as being a receptacle for "desired cells." desirable for developing an receptacle is "artificial organ". The development of the receptacle requires an undesirably long period of time of about 6 10 11 weeks.

Genetically altered or unaltered cells provide a 12 desired metabolic effect. Examples of gene transfer 13 14 technology to produce altered cells are provided in the following three articles: Wolff, et al., "Expression of 15 Retrovirally Transduced Genes in Primary Cultures of 16 Adult Rat Hepatocytes", Proc. Natl. Acad. Sci. USA 84 17 (May 1987): 3344-3348; Ledley, et al., "Retroviral Gene 18 Transfer into Primary Hepatocytes: Implications for 19 Genetic Therapy of Liver-Specific Functions", Proc. Natl. 20 Acad. Sci. USA 84 (1987) 5335-5339; and Wilson, et al., 21 22 "Retrovirus-Mediated Transduction of Adult Hepatocytes", Proc. Natl. Acad. Sci. USA 85 (May 1988) 3014-3018. The 23 art is lacking a satisfactory means to 24 genetically altered or unaltered cells into an organism 25 26 and maintain those cells permanently within that organism 27 that the organism benefits from the desired such

metabolic effect of the cells.

1	The field of angiogenesis has been severly limited
2	by the absence of devices and well defined methods for
3	the selective demonstration of new blood vessel or
4	"neovessel" growth. The importance of site-directing
5	physiological neovessel formation has been long
6	recognized in medicine. The prior art has indicated the
7	possibility of such a process, but does not provide a
В	neovessel design in the form of physiological embodiments
۵	for this purpose

10 The invention is an in vivo site directed 11 neovascularization device. The device includes a 12 support. The support can be an absorbable support, a 13 non-absorbable support, or both. The device also 14 includes a biological response modifier for inducing 15 neovascularization. The biological response modifier is 16 adsorbed to support.

17 The invention also includes a method for directing 18 in vivo neovascularization. The method requires 19 adsorbing a biological response modifier for inducing 20 neovascularization onto a support. The step of 21 contacting a therapeutically effective amount of said 22 adsorbed biological response modifier to at least one 23 selected tissue then occurs. The method then involves 24 directing or culturing neovascular cell growth at the 25 contacted, selected tissue for a sufficient time to 26 obtain a vascular structure.

27 The method of this invention is useful for 28 providing artificial organs.

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Objects of the present invention are to provide:

2	(1) a new device for inducing site-directed
3	neovascularization; (2) a method for in vivo formation of
4	new blood vessel or a vascular bed; (3) mammalian cells
5	collected about the implanted device of the present
6	invention for multiplication, cloning, manipulation and
7	implantation thereof; (4) a vascular bed for
8	transplantation; and (5) other objects made evident from
9	the following detailed description of the invention.
10	BRIEF DESCRIPTION OF THE DRAWINGS
	MALE PROJECT TEST OF THE BUILDING
11	Figure 1 illustrates ECGF binding to collagen
12	supports.
13	Figure 2 illustrates the effect of implanting ECGF
14	immobilized on collagen sponges and the results thereof
15	(arrows to sponges) are shown.
16	Figure 3 illustrates the H & E histological stain
17	of sponges (IP in rat) are shown.
18	Figure 4 illustrates the site-directed gelfoam
19	implant (Sg) with GF (growth factor) between liver (left,
20	L) and spleen (right, Sp).
20	n, and apream (rryme, ap).
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21	Figure 5 illustrates genetically engineered rat

Figure 6 illustrates a cross-section of a blood vessel developed according to this invention.

ECGF at 4 to 6 weeks of post-implantation.

hepatocytes recovered from collagen sponges adsorbed with

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Figure 7 illustrates an angiogeneic response

2	induced by HBGF-1 \underline{in} \underline{situ} four weeks after surgery.
3	Figure 8 illustrates the posterior portion of a
4	fiber implant containing vascular strings that are
5	generally connected to the mesentary tissue around the
6	bowel loop.
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7	Figure 9 illustrates multiple vascular connections
8	between the fiber implant and mesenterial vessels and
9	vascular turbosity within the implant.
10	Figure 10 illustrates an x-ray view of the
11	multiple vascular connections of Figure 9.
12	Figure 11 illustrates a histological examination
13	of a longitudinal section that reveals the presence of
14	multiple vascular lumina surrounded by thick, collagenous
15	and muscular walls of the neovessel structure.

20 . Figure 13 illustrates serum bilirubin levels of a 21 Gunn rat implanted with hepatocytes seeded onto collagen

Figure 12 illustrates the vascular bundle of

Figure 6 at higher magnification which reveals the rich

collagen component of the vascular structure and

abundance of endothelial cell-lined capillary structures.

- 22 (Type IV) and HBGF-1 coated PTFE fibers.
- Figure 14A illustrates a Gortex shunt tube, containing a collagen I (Gelfoam) sponge, impregnated with HBGF-1, implanted onto the aorta of a rat for one
- 26 month, then excised and cross-sectioned.

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Figures 14B, 14C and 14D illustrates a Gortex 1 2 shunt tube containing a bundle of Gortex angel-hair 3 fibers coated with Type I collagen and impregnated with HBGF-1. 4

DETAILED DESCRIPTION OF THE INVENTION

The invention includes both a composition or "device" and a method for using that device. The device vivo to stimulate and is used in neovascularization. The neovascularization 10 accompanied by the growth of other cellular tissue 11 including nerves. The device requires a support. The 12 support must be capable of adsorbing response modifier or adhering to a composition that can 13 adsorb a biological response modifier. 14 The biological 15 response modifier is a compound that stimulates and 16 induces neovascularization. The invention further includes a method for inducing neovascularization that 18 can include the development of artificial organs and/or 19 genetically engineered tissues.

A biological response modifier can be at least one compound or agent that stimulates or facilitates vascular cell growth from a tissue or organ. In other words, a biological response modifier is a biochemical agent, such factor. hormone. or their chimeric as growth derivative, that directly or indirectly induces transcriptional or translational cellular event. A biological response modifier directly or indirectly exerts an effect through a high affinity receptor. This effect produces vascular cell growth. Compounds that stimulation of a receptor include exert direct hormones. Compounds that provide indirect stimulation of

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a receptor include hormone prototypes or precursors and hydrolases. Hydrolases, such as a plasminogen activator, collagenase, or heparinase, initiate a biological response by enzymatically activating or releasing latent, stored, or zymogen precursors of direct biological

stored, or zymogen precursors of direct biological

6 response modifiers.

7 Biological response modifiers desirable angiogenic 8 growth factors include a member of the group consisting 9 of HBGF-I, HBGF-II, platelet-derived growth 10 (PDGF). macrophage-derived growth factor (MDGF). 11 epidermal growth factor (EGF), tumor angiogenesis factor 12 (TAF), endothelial cell growth factor (ECGF), fibroblast 13 growth factor (FGF), hypothalamus-derived growth factor 14 (HDGF), retina-derived growth factor (RDGF), and mixtures 15 thereof. The preferred embodiment of the invention uses 16 HBGF-I. Desirable hydrolases include a member selected 17 from the group consisting of heparinase, collagenase, 18 plasmin, a plasminogen activator, thrombin, heparatinase. 19 and mixtures thereof.

20 Hormones such as the growth factors 21 particularly desirable for use in this invention as 22 biological response modifiers. Hormones specifically 23 elicit cell growth and differentiation. The use of 24 hormones as biological response modifiers 25 neovascularization to rapidly occur and to form a 26 complete vascular structure. The resulting blood vessel 27 stimulated by hormones is more than just a mass of cells 28 in that it has a tubular cavity and connective tissue between its cells. The resulting blood vessel produced 29

from the use of hormones is complete within itself and
can be excised and transplanted into another portion of
the body. The other biological response modifiers
produce similar results, but do not provide as rapid a
growth as hormones and, in particular, the HBGF-I and
HBGF-II hormones.

7 The invention includes a biocompatible support to 8 which the biological response modifier is adsorbed. 9 The support can be either or both an absorbable or 10 non-absorbable biocompatible matrix. The support must be 11 implantable into an organism and is, desirably, rigid and 12 strong enough to be transplantable 13 neovascularization has occurred. The biocompatible 14 support must have the rigidity and strength to support 15 Examples of absorbable supports neovascularization. 16 include a member selected from the group consisting of 17 collagen Type I, known commercially by the trade name laminins, 18 "Gelfoam", fibronectins. gelatins. 19 glycosaminoglycans, glycolipids. proteolipids. 20 mucopolysaccharides, glycoproteins, polypeptides, 21 mixtures thereof. Examples of non-absorbable matrices 22 include members of the group consisting of nylon, rayon, 23~ polypropylene. polyethylene, expanded PTFE. 24 cross-linked collagen Type IV, and mixtures thereof. 25 desirable that a selected support contain 26 extracellular matrix protein to provide or to facilitate 27 the adsorption of the biological response modifier to the 28 biocompatible support.

1 An extracellular matrix protein can be 2 material from which the biocompatible support is formed 3 or a component added to the biocompatible support to 4 provide or, alternatively, facilitate 5 adsorption of the biological response modifier to the 6 biocompatible support. An extracellular matrix protein 7 component can include a pure or mixed composition of 8 proteins or polypeptides. The proteins and polypeptides 9 can be either natural or synthetic. The extracellular 10 protein component is desirably derived from 11 extracellular structural molecules. These extracellular 12 structural molecules include a member selected from the 13 group consisting of collagens, laminins, fibronectins, 14 gelatins, glycosaminoglycans, glycoproteins, 15 proteoglycans, and mixtures thereof.

16 Expanded polytetrafluoroethylene (PTFE) has been 17 found to be most suitable non-absorbable support for this 18 invention. This support provides the 19 PTFE has a general lack of an inflammatory benefits. 20 response which is the basis for the current acceptance of PTFE in the surgical community. PTFE can be coated 21 22 conveniently with various components of the

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1 cellular matrix which can adsorb a biological response 2 modifier. Biologically active HBGF-1 and HBGF-2 can be 3 immobilized to collagen-coated PTFE by previously 4 established methods. PTFE polymers are 5 engineered to various specifications to meet a multitude 6 of required configurations.

7 The configuration of the non-absorbable PFTE is a 8 more critical aspect of the long-term implant model. All g multicellular organisms utilize a three-dimensional 10 architecture of branching fiber networks to solve the 11 problem of increasing surface area in a given volume. 12 Seeding of such a network with HBGF polypeptides before 13 implantation allows for high localized concentrations of 14 the mitogen. Non-woven multifilament angel-hair fibers 15 of expanded PTFE are commercially available from W.L. 16 Gore and Associates, Inc., Flagstaff, Arizona. These 17 fibers allow sufficient organized surface 18 infiltrating cells to be exposed to the environment of 19 the host. This permits the free exchange of nutrients 20 and toxic waste to occur while neovascularization 21 processes occur. Furthermore, cell shape as determined 22 by cytoskeletal components and attachment to a specific 23 matrix generally is regarded to play a significant role 24 in both cell proliferation and differentiation.

A support can be provided for use in this invention in any desired shape and size. A support as small as one 1mm² is suitable to provide a base for neovascularization. Desirable shapes for a support can

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1.	be a strip, a sponge, or a tube. Supports are desirably
2	capable of being secured within an organism. Suitable
3	means for securing a support can include a staple,
4	biocompatible glue, or other surgical procedures such as
5	suturing or tying the support to a tissue.

A desirable support is obtained by filling a tube or sleeve of expanded PTFE with expanded PTFE fibers or "angel hair". Supports formed from tubes or sleeves provide a pouch for an artificial organ. The tubular form of the support and the bundle of fibers within the tube are particularly desirable for directing neovascularization. Such embodiments can be receptacles for implanted cells when the invention is used to provide an artificial organ.

15 The most effective concentrations for a biological 16 response modifier can be any concentration that elicits a 17 growth response from the target cells, but is not toxic 18 to those cells. Effective or therapeutic concentrations 19 of angiogenetic growth factors are between about 1 to 20 about 10 nanograms per cubic millimeter of a support. 21 support for this calculation includes both the absorbable 22 support and the non-absorbable support.

A support is provided in an amount suitable to
establish the length and width of the desired blood
vessel. For example, if a blood vessel is desired between
two tissues and there exists a distance between those two
tissues, then a corresponding length of support is

2 length and width of this desired blood vessel. The 3 amount of the biological response modifier is then 4 adapted to the amount of support required to form this

implanted into the organism to provide the approximate

5 basic structure.

6 The invention can be practiced without a 7 non-absorbable support. For example, a complex with 8 HBGF-1, or HBGF-2 is capable of inducing 9 neovascularization in vivo at polypeptide concentrations 10 consistent with the demonstration of this biological 11 activity in vivo. This neovascular response is capable 12 of sustaining induced site-specific neovessel formation 13 for up to four weeks in the neck and peritoneal cavity of 14 the rat. However, the device of this invention without a 15 support has limited utility the for induction 16 long-term neovessels. This is because the 17 three-dimensional architecture of the collagen sponges is 18 ultimately disrupted by a reabsorption process that 19 occurs within three to four weeks after implantation. 20 Nonabsorbable solid polymeric supports of well-defined 21 specifications and containing bonded components 22 extracellular matrices induced the expression of 23 long-term stable neovessels in vivo. An example of such 24 an embodiment is a nonabsorbable support bonded with both 25 collagens Type I and Type IV and having both HBGF-1 and 26 HBGF-2 attached to the collagens.

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A neovascularization device can also be seeded 1 2 with desired cells prior to or subsequent to implantation 3 in a host. In a preferred embodiment, such cells are 4 mammalian cells and express a protein capable 5 performing a particular function. The cells can be 6 genetically engineered cells capable of expressing a 7 heterologous protein. Alternatively, the cells can be 8 naturally occurring cells capable of providing a desired function or functions such as hepatocytes.

Desirable embodiments of the invention have cells
seeded in or on the neovascularization device which are
genetically engineered to express at least one
heterologous protein. Such a protein is preferably a
therapeutic agent. The expressed protein may or may not
be secreted from the genetically engineered cells.

16 The genetically engineered cells used with this 17 invention are transformed with at least one gene that 18 encodes for the desired heterologous protein. The cells 19 are transformed with a suitable vector or expression 20 vehicle which includes the desired gene. The vector can 21 also include a promoter for expression in the host 22 cells. In mammalian cells, the promotor for expression can be SV 40, LTR, metallothionein, PGK, CMV, ADA, TK, or 23 24 others. The vector can also include a suitable signal 25 sequence or sequences for secreting the therapeutic agent 26 from the cells. The selection of a suitable promotor is

deemed to be within the skill of the art.

The vector or expression vehicle is preferably a 1 viral vector and in particular a retroviral vector. 2 3 Representative examples of suitable viral vectors, which can be modified to include a gene for a therapeutic 4 agent, include Harvey Sarcoma virus, ROUS Sarcoma virus, 5 6 MPSV, Moloney murine leukemia virus, DNA viruses such as 7 adenovirus and others. Alternatively, the expression 8 vehicle can be a plasmid. Transformation 9 accomplished by liposome fusion, calcium phosphate or transfection. electroporation. 10 dextran sulfate lipofection, tungsten particles, or other procedures. 11 The selection of a suitable vehicle for transformation is 12 deemed to be within the scope of those skilled in the 13 14 art.

When a retroviral vector is employed as the 15 16 expression vehicle for transforming cells, steps should 17 be taken to eliminate and/or minimize the chances for replication of the virus. Various procedures are known 18 in the art for providing helper cells which produce viral 19 20 vector particles that are essentially free of replicating 21 virus. Examples of such procedures are found Markowitz, et al., "A Safe Packaging Line for Gene 22 23 Transfer; Separating Viral Genes on Two Different 24 Plasmids". Journal of Virology 62(4) (April 1988):1120-1124; Watanabe, et al., "Construction of a 25 Helper Cell Line for Avian Reticuloendotheliosis Virus 26 Cloning Vectors", Molecular and Cellular Biology 3(12) 27 28 (Dec. 1983):2241-2249; Danos, et al., "Safe and Efficient 29 Generation of Recombinant Retroviruses with Amphotropic

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1 and Ecotropic Host Range", Proc. Natl. Acad. Sci. 85 2 (Sept. 1988):6460-6464; and Bosselman. 3 "Replication-Defective Chimeric Helper Proviruses and 4 Factors Affecting Generation of Competent Virus; 5 Expression of Moloney Murine Leukemia Virus Structural Genes via the Metallothionein Promoter", Molecular and 6 7 Cellular Biology (5) (May 1987):1797-1806 disclose 8 procedures for producing a helper cell which minimizes g the chances for producing a viral particle that includes 10 replicating virus. This procedure and other procedures can be employed for genetically engineering cells by use 11 12 of a retroviral vector. In addition to the promotor and 13 the gene for the therapeutic agent, other material can be 14 included in the vector. This material can include a 15 selection gene such as a neomycin resistance gene, a 16 sequence for enhancing expression, or other materials.

17 Genetically engineered mammalian cells can be 18 implanted in a mammal by use of a neovascularization 19 device. These genetically engineered cells are desirably 20 implanted into a mammal of the same species. In a 21 preferred embodiment, the genetically engineered 22 mammalian cells are cells originally derived from a 23 patient, genetically engineered to include a gene for at 24 least one therapeutic agent, and implanted into the 25 patient from which they were derived by use of a 26 neovascularization device in accordance 27 invention. These autologous genetically engineered cells 28 then provide "gene therapy" by in vivo production of the 29 therapeutic agent for treatment of the patient.

1 The genetically engineered cells can be engineered 2 such that the therapeutic agent is secreted from the 3 cells in order to exert its effect upon cells and tissues 4 either in the immediate vicinity or in more distal 5 locations. Alternatively, the therapeutic agent, if it 6 is not secreted from the engineered cells, exerts its 7 effect within or on the engineered cells and can cause 8 the metabolism of substances that diffuse into or onto the cells. Examples of such therapeutic agents include 9 10 adenosine deaminase (ADA) that functions within the cell to inactivate adenosine, 11 a toxic metabolite 12 accumulates in severe combined immunodeficiency syndrome, 13 or phenylalanine hydroxylase that functions within a cell 14 inactivate phenylalanine, a toxic metabolite in 15 phenylketonuria.

16 The genetically engineered cells used with this 17 invention are transformed with a gene for at least one 18 heterologous protein. This protein is preferably a 19 therapeutic agent. The term "therapeutic agent" is used 20 in its broadest sense and means any agent or material which has a desired or beneficial effect on the host. 21 22 The therapeutic agent can be more than one type of 23 Desirable proteins include CD-4. Factor VIII. 24 Factor IX, von Willebrand Factor. TPA. urokinase. 25 hirudin, the interferons, tumor necrosis factor, the 26 interleukins, hemotopoietic growth factors 27 G-CSF. GM-CSF. IL3, erythropoietin, antibodies,

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skilled in the art.

1 glucocerebrosidase, ADA, phenylalanine hydroxylase, human 2 growth hormone, insulin and others. The selection of a 3 suitable gene is deemed to be within the scope of those 4 skilled in the art. Mixtures of cell types can also be 5 used with this invention such s genetically engineered 6 smooth muscle cells. fibroblasts. glial cells. keratinocytes, or others.

The effect in genetically engineered cells when used in gene therapy, can be controlled by the selection of high producing clonal populations and/or the use of vectors with enhanced expression. This can provide, in vivo, therapeutically effective amounts of a desired therapeutic agent for treating a patient. In determining the number of cells to be implanted, factors such as the half life of the therapeutic agent, volume of the vascular system, production rate of the therapeutic agent by cells, and the desired dosage level are considered. The selection of such vectors and cells is dependent on the therapeutic agent and is within the scope of those

21 The neovascularization device of the invention can 22 also be employed to obtain cells from a host by 23 implanting the device in a host and after a period of 24 time removing the implanted neovascularization device 25 from the host for recovery of cells which have been 26 collected on the device. Such cells can be 27 differentiated and used for a variety of purposes. For

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example, this procedure can provide a source of autologous cells for genetic engineering and subsequent return to the host as genetically engineered cells for expression of a protein. Cells collected in this manner can be genetically engineered and then returned to the host to provide an artificial organ.

The process for directing neovascularization first involves preparing the device of this invention as described above. The device is prepared by adsorbing a biological response modifier, that is suitable for inducing neovascularization, onto support. а biological response modifier must be present on the support in such a concentration as to be therapeutically effective for eliciting cell growth. The adsorbed biological response modifier is then contacted to at least one selected tissue. Typically, the device is connected to at least two separate sites between which a blood vessel is desired. These two sites can be the same or separate tissues or organs. The method then involves culturing neovascular cell growth at or from the contacted tissue. Culturing of the contacted cells must for a sufficient time to allow or enable neovascularization and the vascular structure to form.

Figure 1 demonstrates that ECGF binds to collagen supports. This is shown by an elution profile of HBGF-1 (ECGF) from collagen type IV-Sepharose and gelatin-Sepharose columns. Collagen Type IV-Sepharose and The gelatin-Sepharose (1 ml) were packed in a column

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1 and washed with 5 mls of 2M NaCl in 50mM Tris HCl, pH 2 7.4, followed by an exhaustive wash with 50mM Tris HCl, 3 pH 7.4 (adsorbtion buffer; AB). The Gelatin-Sepharose 4 was from Pharmacia. Bovine collagen-Type IV-Sepharose 5 was obtained from Sigma Chemical Company, St. Louis, MO. 6 and (125I)-HBGF-1 was prepared as previously described. 7 (1251)-HBGF-1 (approximately 5X105 cpm) in absorption 8 buffer was added to the column in volume 9 the column washed with approximately 0.1 ml and 10 absorption buffer. Elution column-associated of 11 (125I)-HBGF-I was achieved with 1.5M NaCl in absorption 12 buffer or 50 units of heparin (Upjohn, Kalamazoo, MI) in 13 absorbtion buffer. The NaC1-eluted column was 14 regenerated with an absorption buffer wash and the 15 heparin-eluted column was regenerated by consecutive 16 washes with 1.5M NaCl in absorption buffer followed by 17 another wash with absorbtion buffer. The matrix affinity 18 procedures were performed at room temperature (about 22°C 19 to 25°C).

Figure 2 demonstrates that ECGF binds to collagen supports. The adsorbed factor was implanted in various anatomical sites to demonstrate the practicality of using growth factor-adsorbed implants to stimulate neovessel formation and the growth of vascular beds in areas of interest. The effect of implanting ECGF immobilized on collagen sponges and the results thereof (arrows to sponges) are shown:

1	A. Neck, 2 weeks, no ECGF;
2	B. Neck, 2 weeks, plus ECGF;
3	C. IP, 2 weeks, no ECGF;
4	D. IP, 2 weeks, plus ECGF;
5	E. IP, 2 weeks, plus ECGF site-directed; and
6	F. IP, 2 weeks, plus ECGF implantation in
7	omentum.
8	Figure 3 demonstrates that the device of this
9	invention induces significant angiogenesis in situ.
10	These implants were removed at various times for
11	examination by common methods of histology in order to
12	determine the microscopic nature of these dynamics. The
13	following abbreviations are used: Sg represents "sponge
14	(C-1)"; Sp represents "spleen"; L represents "liver"; and
15	BV represents "blood vessel (aorta)". H & E histological
16	stain of sponges (IP in rat) are shown:
17	A. spongetwo weeks, IP, without ECGF;
18	B. spongeone week, IP, plus ECGF;
19	C. spongetwo weeks, IP, plus ECGF;
20	D. sponge glued to liver, 2 weeks, plus ECGF;
21	E. sponge glued to spleen, 2 weeks, plus ECGF;
22	and
23	F. sponge wrapped around aorta, 2 weeks, plus
24	ECGF.

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Figure 4 demonstrates that ECGF induces significant and stable angiogenic response in situ by the recruitment of appropriate cell types as shown in Figures 2 and 3. Implants were established to create site-directed bridges between a large variety of organs, vessels, tissues and the like. Illustrated are the site-directed Gelfoam implant (Sg) with growth factor (GF) between liver (Left, L) and spleen (right, Sp).

Figure 5 demonstrates that the device of this invention serves to create neovessels independent of the implantation site in situ. The device has an ability to serve as a recruitment vehicle for mammalian cells in general and as a vehicle to maintain the viability and physiological environment for and of the implanted and transplanted cells. Genetically engineered rat hepatocytes recovered from collagen sponges adsorbed with ECGF after 4 to 6 weeks post-implantation are shown. Hepatocytes were removed to determine their viability.

Figure 5A shows the results with no growth factor. Note that in Figure 5A few cells appear to be unhealthy and there is no proliferation or growth of survivor cells. Figure 5B shows the results with growth factor. Note that in Figure 5B healthy viable cells are accompanied by significant proliferation.

The device and method of this invention can provide angiogenesis and neovascularization from one or more sites on a single tissue or different tissues. The development of a blood vessel from a single site of one

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tissue, such as an artery, provides a vessel that can be transplanted or that can be used as an artificial organ.

The development of a blood vessel between two or more sites located on the same or different tissues provides improved circulation between the sites.

Figure 6 illustrates a cross section of a blood vascular structure developed by the device and method of this invention. This figure demonstrates that the blood vessels developed by this invention are not merely a bundle of vascular cells growing in an manner. The blood vessel 1 contains endothelial cells 2, mesothelial cells 3, pericytes 4, smooth muscle cells 5, fibroblasts 6, and neuronal-like cells 7. The cross section of the blood vessel 1 demonstrates the formation of capillary-like structures 8, arteries 9, and vein-like structures 10. This development of a complete vascular structure provides а rigid vessel that remains permanently in the organism and that can be transplanted within this organism.

A method of this invention can be used to provide an artificial organ by first directing the growth and development of a blood vessel from a tissue. The developed blood vessel is then injected or seeded with cells from a selected tissue or organ. The injected cells can be genetically altered before being seeded into the blood vessel. The seeded cells can provide a desired metabolic effect. These metabolic effects can include

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1 hepatic functions such as bilirubin metabolism and pancreatic functions such as insulin production. 2 metabolic functions can be provided by cells containing 3 one or more hormone producing genes. Artificial organs 4 developed according to this invention can provide desired 5 б functions without being subject to a response from the 7 organism's immune system.

EXAMPLE 1

Example 1 demonstrates various embodiments of the device or composition of the invention and the method by which the device is produced. This example uses HBGF-I with a radioactive iodine marker. In therapeutic use, the radioactive marker would not be present. Example 1 is as follows.

15 Gelatin-Sepharose and collagen Type IV-Sepharose were examined for the ability to absorb (1251)-HBGF-1. 16 Figures 1C and G show that the majority or approximately 17 80 percent of the (1251)-HBGF-1 binds to immobilized 18 gelatin and collagen Type IV and can be eluted with 1.5M 19 Adsorbed (1251)-HBGF-1 20 can also be eluted 21 with 0.5M NaCl (data not shown). Denaturation of 22 (125_I)-HBGF-1 by heating at 90°C for 23 significantly reduces the ability of the polypeptide to 24 bind 'to immobilized gelatin and collagen Type IV by inactivation of the binding domain within the HBGF-1 25 polypeptide structure. 26

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1	The (1251)-HBGF-1 adsorbed to immobilized gelatin
2	and collagen Type IV can also be eluted with heparin as
3	shown in Figures 1A and E. Approximately 20% of the
4	growth factor, which remains bound after heparin elution,
5	can be eluted with 1.5M NaCl.
б	Pretreatment of the gelatin and collagen Type IV
7	matrix with 50 units of heparin significantly reduces the
В	ability of either matrix to absorb (1251)-HBGF-1 as shown
9	in Figures 1B and F. Regeneration of either matrix with
10	a 1.5M NaCl wash permits (1251)-HBGF-1 adsorption.

Bovine serum albumin at lmg per ml and human tilm fibronectin at lmg per ml do not significantly elute (1251)-HBGF-1 absorbed to either matrix as shown in Figures 1D and H.

15 EXAMPLE 2

Example 2 demonstrates the method for implantation of the device of this invention and for eliciting neovascularization. The use of immobilized gelatin with HBGF-I represents the preferred embodiment of the invented method. Example 2 is as follows.

Example 2 demonstrates that HBGF-I binds to both immobilized gelatin and to collagen Type IV. It is shown that HBGF-I, adsorbed to gelatin sponges, promotes

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angiogenesis in the rat at concentrations of the growth factor which are consistent with the growth factor's activity as an endothelial cell mitogen in vitro. This concentration is about 10⁻³ times lower than the concentration used in vitro in the art.

The abdomen of an anesthesized male rats weighing 250 grams was washed with 20 percent volume to weight (v/w) ethanol and an incision was made into the abdominal cavity wall to expose the abdominal cavity. manufactured by Upjohn, Kalamazoo, Michigan, was cut into strips of approximately 5 by 20mm. The sponge was cemented to the distal area of the abdominal aorta with n-butylcyanoacrylate. A bridge was created with the free end of the sponge when the free end was cemented to another tissue. In the studies that were conducted to provide these examples. the following tissues were actually contacted by the device. These tissues were other organs including the liver, kidney, and spleen, the abdominal cavity, and other macro and micro vessels. Various concentrations of HBGF-1 from about 1 to about 10 ng per mm³ were adsorbed to sponges for these studies. The surgical opening was closed with a staple gun. animals were fed a normal diet and the incision was opened 1 week after surgery. The collagen sponge was surgically extracted, grossly examined for blood vessel formation and the sponge prepared for histological examination.

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It is known that HBGF-1 binds to immobilized gelatin and collagen Type IV, therefore, the possibility was evaluated that commercial gelatin sponges sold by the tradename "Gelfoam" adsorbed with HBGF-1 could be utilized as a method for inducing angiogensis in situ. Survival surgery was performed on the rat in order to implant gelatin sponges which were treated with HBGF-1. HBGF-1-adsorbed Gelfoam was independently placed in the neck and peritoneal cavities in the rat. A significant angiogenic response was observed in situ one week after surgery with lng HBGF-1 per mm² (Figure 2). vessels, which migrated away from the tissue site of implantation, were observed macroscopically exclusively within the gelatin sponge. Control sponges without HBGF-1 and sponges adsorbed with HBGF-1 and heparin did not induce neovascularization after one week in vivo. The latter is consistent with the ability of heparin to prevent HBGF-1 adsorption to immobilized gelatin and collagen-Type IV. A titration curve with various concentrations of HBGF-1 was performed using this procedure and results similar to Figure 1 was observed with 1 to 10ng HBGF-1 per mm3 of sponge (data not Histological examination (Figure 3) of the sponge removed after one week in situ revealed new blood vessel growth within the sponge.

Since HBGF-1-adsorbed Gelfoam alone (without more) is an efficient inducer of angiogenesis from the serosa. The ability of immobilized HBGF-1-adsorbed implants to induce and sustain the process of neovascularization within the peritoneal cavity was assessed. Separate

1 surgical implants were cemented as strips of Gelfoam to 2 the abdominal aorta in the rat creating a bridge between 3 this site and either the kidney, spleen, liver, or 4 abdominal wall (Figure 4). After two weeks in vivo, the 5 implants were examined for the extent of angiogenesis. 6 Bidirectional formation of new blood vessels along the 7 HBGF-1-adsorbed gelatin sponge from the liver and aorta 8 was observed. Similar bidirectional results were 9 observed with implants cemented from the aorta to either 10 the kidney, spleen, or abdominal wall (data not shown). 11 Histological examination of these implants yielded 12 results identical to those observed in Figure 3.

13 Induced neovascularization within the peritoneal 14 cavity was also shown to sustain the proliferative 15 potential of a genetically engineered rat hepatocyte cell 16 strain simultaneously implanted with the HBGF-1-adsorbed 17 Gelfoam (Figure 5). Hepatocytes were grown to high 18 density (108 cells) on a Gelfoam sponge. 19 surgical implantation, 10ng of HBGF-1 per mm3 of sponge 20 was added. Control sponges did not contain any adsorbed 21 HBGF-1. Separate surgical implants were cemented as a 22 bridge between the liver and the spleen and allowed to 23 remain in situ for four to six weeks. At this time, the implants were removed, digested with either trypsin or 24 25 collagenase to recover implanted cells which 26 maintained in tissue culture. Cells which were recovered from HBGF-1-adsorbed Gelfoam sponges 27 were able 28 proliferate in vitro under selective pressure which

reflected genetic disposition (Figure 5B). In contrast, the cells recovered from control Gelfoam sponges displayed a loss of proliferative potential (Figure 5A). Histological examination of sponges containing the cells revealed that HBGF-1 also induced a response similar to Figures 3 and 4.

In accordance with the device and method of the present invention, angiogenesis and neovascularization has been achieved between various tissues and organs as demonstrated by Figures 2 through 5. Neovascularization has been similarly accomplished between the following loci (data not shown): liver to spleen; liver to kidney; spleen to kidney; liver to acrta; liver to vena cava; liver to omentum (omentum, containing pancreatic tissue); acrta/to vena cava; spleen to omentum kidney to acrta; kidney to vena cava; spleen to omentum; omentum to acrta; and omentum to vena cava;

EXAMPLE 3 AND COMPARATIVE EXAMPLE A

Example 3 demonstrates the device of the invention
having a non-absorbable support. The experiments
performed to derive this example were conducted with
either Type I or Type IV collagen and involved
implantation onto the liver or the spleen of a rat.

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Comparative Example A demonstrates that the use of the same materials and procedures of Example 3 without HBGF-1 did not induce neovascularization.

HBGF-1 adsorbed, collagen-coated (Type I or IV) expanded PTFE fibers were surgically implanted in the peritoneal cavity (onto the liver or the spleen) of the A significant angiogenic response was specifically induced by HBGF-1 in situ and the results four weeks after surgery are shown in Figure 7. Blood vessels, which have migrated from the tissue site of implantation, could be observed macroscopically within and around the implanted fibers. The anterior portion of the fiber implant, which was attached to the liver, exhibited substantial neovessel growth from the liver into the interior of the implant (Figure 7). Further examination revealed that the posterior portion of the fiber implant (attached to a specific organ) or regions in the vicinity of the implant contained vascular "strings" which were generally connected to the mesentary tissue around the bowel loop (Figure 8). It was also possible to induce and sustain long-term bi-directional neovessel formation between the liver and spleen by the implantation of separate HBGF-1-treated fibers on each organ. The ability of HBGF-1 adsorbed implants to maintain the neovessel structures within the peritoneum is evidenced by these highly vascular bridges. Control fibers of Comparative Example A did not induce neovascularization even after six months following surgical implantation. Titrations with various concentrations of HBGF-1 were performed using this procedure. Similar results were

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1	obtained with HBGF-1 at concentrations between 1 to 100
2	ng/mm3 of fiber surface area. The concentration of
3	HBGF-1 required to induce an angiogenic response in the
4	fiber implant model is consistent with the results
5	obtained with the Gelfoam implant model and the mitogenic
6	activity of the polypeptide in vitro.

EXAMPLE 4

Example 4 demonstrates that the blood vessel produced in Example 3 displayed a large organized solid matrix including a network of neovessel formations.

11 Two months following surgical placement of the 12 HBGF-1-treated implant on the spleen of a rat, the 13 abdominal organs were perfused and fixed (formaline) using a catheter placed in the lower thoracic aorta. 14 15 Subsequently, the abdominal organs were perfused with a 16 radio-opaque silicone rubber dye sold by the trademark, 17 Microfil, followed by soft X-ray analysis (magnification 18 Multiple vascular connections between the fiber 19 implant and mesenterial vessels were observed as well as 20 a vascular turbosity within the implant which is typical 21 for new vessel formation (Figure 9). 22 examination of the implant itself displayed a large 23 organized solid matrix containing a network of neovessel 24 formations interdigitated with different cell types, 25 which is consistent with results previously obtained with 26 the short-term HBGF-1-treated Gelfoam implant model. 27 X-ray analysis of the long-term fiber implant as shown in 28 Figure 10 has confirmed that neovessel formation within 29 the fiber network has become integrated with the vascular 30 tree of the host, primarily through the bridges

1 ("strings") of richly vascular tissue (Figures 7 and 8). 2 Histological examination of the longitudinal section 3 through a typical vascular connection revealed 4 presence of multiple vascular lumina surrounded by thick, 5 collagenous and muscular walls of the neovessel structure 6 (Figure 11). Cross-sectional analysis through these 7 vascular connections further related the presence of a 8 of mesothelial cells surrounding a large 9 vascular lumina in the central portion, encompassed by 10 prominent endothelial cells and multiple layers of smooth 11 muscle cells. representing mature and 12 differentiated arteries. Venous lumina are less visible 13 and present as partially collapsed slits. Within the 14 periphery are abundant capillary lumina, and the entire 15 surrounded vascular bundle is by 16 fibrocellular capsule (Figure 6). Further examination of 17 this resource at higher magnification revealed the 18 relatively rich collagen component of vascular structure 19 as well as the abundance of endothelial cell-lined 20 capillary structures (Figure 12). The presence of two 21 distinct, yet prominent, round structures, marked with 22 asteriks were also observed. These structures displayed 23 histological characteristics o£ neuronal-like 24 structures. Collectively these data suggest that HBGF-1 25 is capable of signaling a variety of the squamous 26 mesothelial cells of the serosa and the proximal cells of 27 the tunica adventita to initiate angiogenesis. 28 appearance of mesoderm- and neuroectoderm-derived cells 29 is consistent with the ability of HBGF-1 to act as a 30 mitogen in vitro for epithelial cells, fibroblasts, 31 smooth muscle cells, mesothelial cells. endothelial

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- cells, astrocytes and oligodendrocytes. The presence of 2 neuronal-like structures is also consistent with the 3 nerve growth factor (NGF)-like biological activity of 4 HBGF-1 to induce neurite extension and survival of PC12 5 cells in vitro.

EXAMPLE 5 AND COMPARATIVE EXAMPLE B

Example 5 demonstrates that the presence of a 8 large organized solid matrix, containing a network of 9 mature muscular neovessel formations of Example 4 and 10 which are contiguous with the host's vascular tree in 11 situ, permits successful selective cell transplantation.

12 Comparative Example B demonstrates that the use of 13 the same materials and procedures of Example 5 without 14 HBGF-1 did not sustain selective cell transplantation.

15 Homozygous Gunn rats lack 16 UDP-glucuronosyltransferase for bilirubin and cannot 17 efficiently excrete bilirubin. For this reason, Gunn 18 rats exhibit lifelong nonhemolytic unconjugated 19 hyperbilirubinemia. In order to examine the genetic 20 therapy potential of this system, hepatocytes 21 harvested by collagenase perfusion of syngeneic Wistar 22 (RHA) rats. The Wistar rat is genetically identical to 23 the Gunn rat except that it contains a normal bilirubin 24 conjugation locus.

In Example 5, HBGF-1 adsorbed collagen (Type IV) coated PTFE fibers were implanted next to the liver and after ten to fourteen days the peritoneal cavity was

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1 surgically opened revealing numerous neovessel formations 2 both protruding from the liver and extending into the 3 bundle of fibers (Figure 7) and connecting the bowl loop 4 with richly vascular bridges. Primary 5 harvested from syngeneic Wistar (RHA) rats were injected 6 into the fiber network of the vascularized fibers. 7 Immediately, serum bilirubin levels began to decrease and 8 ten days after hepatocyte injections, the serum bilirubin 9 levels had decreased by 50 percent. A gradual decrease 10 to greater than 60 percent was observed for the duration 11 of the experiment (60 days) as shown in Figure 13A. 12 Experiments have determined that reduced levels of serum 13 bilirurin (>60%) can be maintained at least 181 days and histological examination of these long-term implants 14 15 contain viable hepatocytes. These data suggest that 16 HBGF-1 fiber implant model functions in vivo as a 17 receptacle for the successful site-specific introduction 18 of cells capable of expressing differentiated а 19 physiologic function.

In Comparative Example B, the hepatocytes were seeded onto collagen (Type IV) coated PTFE fibers, which did not contain adsorbed HBGF-1, and surgically implanted on the right lobe of the liver. The serum bilirubin levels decreased to approximately 50 percent. This was followed immediately by a sharp reversion to the original serum bilirulrin level. Figure 13B shows that the serum bilirubin levels remained constant for the duration of the experiment (60 days). Histological examination of these implants after twenty davs suggested accumulating levels of toxic-like acids within the fiber implant led to the ultimate death of the transplanted hepatocytes.

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The long-term HBGF-1 fiber implant model of Example 5 induces a prominent angiotropic and neurotropic when appropriately implanted in the rat. Example 5 demonstrates the ability of HBGF-1 to induce, sustain, and maintain the anatomical coordination of highly sophisticated and widely diversified mammalian cell types in vivo. The interrelationships between extracellular matrix components differentiation-specific gene regulation can provide information critical for genetic engineering therapies. 10 11 This invention may also prove useful as a site-specific transgenic alternative with the ability to understand the 12 13 and coordinated expression of growth and temporal 14 differentiation signals during neuronal and angiogenic 15 development in the adult.

16 EXAMPLE 6

Example 6 demonstrates the neovascular device of 17 18 this invention wherein genetically engineered cells are 19 seeded into the device. Example 6 is as follows.

The construction of the pG2N retroviral vector, that was used to genetically engineer endothelial cells to produce rat growth hormone, was performed with SV40 promoted neomycin resistance gene and a rat growth hormone cDNA. These were placed into the pB2 retroviral vector provided by the Laboratory of Molecular Hematology at NIH. A growth hormone cDNA was obtained by digesting the plasmid RGH-1 according to Nature 270 (1977):494 with

1	Xho I and Mae II restriction endonucleases from
2	Boehringer Mannheim Biochemicals. This rat growth
3	hormone cDNA was electrophoretically isolated out of ar
4	agarose gel and purified via binding/elution to glass
5	beads sold by the tradename, Geneclean Bio, 101, La
6	Jolla, California. This growth hormone cDNA was then
7	blunted using the large fragment of DNA polymerase Klenow
8	known by the name, from New England Biolabs and
9	nucleotide triphosphates as recommended by the
10	manufacturer. This fragment was then purified with
11	Geneclean product.

The B2 vector was constructed in order to replace the Neo^R gene in N2 according to M.A. Eglitis, et al., <u>Science</u> 230 (1985):1395; D. Armentano, et al., <u>J. Virol</u> 61 (1987):1647 with a multiple cloning site. N2 was first digested with Eco RI, thereby releasing both the 5' and 3' LTRs with the adjoining MoMLV flanking sequences. The 3' LTR fragment was ligated into the EcoRI site of the plasmid GEM4 from Promega Biotech. The 5' LTR fragment with its flanking gag sequence was then digested with Cla I, Hind III linkers were added, and the fragment was inserted into the Hind III site of poEM4.

The pB2 vector was digested with the HincII restriction endonuclease from New England biolabs, and phosphatased using calf alkaline phosphatase from Bochringer Mannheim Biochemicals. The pB2 plasmid was then purified with the Geneclean product. The pB2 vector and the rat growth hormone cDNA were then ligated using T4 ligase from New England Biolabs, pG2 was then digested

1 with BamHI from New England Biolabs, purified with the 2 Geneclean Bio 101 product, and blunt ended with the 3 Klenow fragment. A 340 base pair SV40 promoted neomycin 4 resistance gene fragment was isolated from the pSV2CAT 5 plasmid (ATCC accession number 37155) by digesting with 6 and HindIII from New England Biolabs. This 7 fragment was isolated by agarose gel electrophoresis and 8 purified with the Geneclean product. The SV40-neomycin 9 resistance fragment was then ligated using T4 ligase from 10 New England Biolabs with pG2 and transformed into DH5 11 competent bacteria per the manufacturer's instructions 12 (BRL). Colonies were screened and the resulting plasmid 13 construct was called pG2N. The SAX vector was obtained 14 described in Proc. Natl. Acad. Sci. USA 83 15 (1988):6563.

16 The recombinant vectors, N2, SAX, G2N, used in 17 this example were each separately transfected into the 18 currently available retroviral vector packaging cell 19 lines, including the amphotropic packaging lines, PA317 20 Mol. Cell. Biol. 6(1986):2895, and the ecotropic line. Psi2, Cell 33(1983):153. These lines were developed in 21 22 order to allow the production of helper virus-free 23 retroviral vector particles.

B. The CD4 containing plasmid, p4B, which was a gift of Richard Axel of College of Physicians and Surgeons Columbia University, New York, New York, was digested with the restriction endonucleases Eco RI and Bam HI from New England Biolabs, Beverly, Massachusetts, to release the CD4 gene which was isolated by agarose gel

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1 electrophoresis followed by purification via 2 binding/elution to glass beads using the 3 product, Bio 101, La Jolla, California, in the manner 4. recommended by the manufacturer. The CD4 fragment was ligated, using T4 DNA ligase as recommended by the 5 6 supplier, into Eco RI plus Bam HI cut Bluescript cloning 7 vector from Stratagene Co., La Jolla, California. 8 ligation was then transformed into competent DH5 alpha 9 bacteria from Bethesda Research Labs, Gaithersburg, 10 Maryland, and white colonies were isolated and screened 11 for proper insert size to yield the plasmid pCDW. To 12 produce a suitable plasmid based expression vector for 13 the CD4 gene, the plasmid SV2neo, obtained from American 14 Type Culture Collection, Rockville. Maryland, 15 digested with Hind III plus Hpa I. A synthetic 16 polylinker sequence from pUC-13 the vector 17 Pharamicia, Piscataway, New Jersey, was inserted via T4 18 DNA ligase in place of the NeoR gene of PsV2neo. 19 ligation was transformed into DH5 bacteria from Bethesda 20 Research Labs and colonies screened for the presence of 21 restriction enzyme sites unique to the polylinker to 22 yield the vector pSVPL. The pSCPL expression vector was 23 further modified by the insertion of an Xho I linker 24 using conditions and reagents suggested and supplied by 25 New England Biolabs, into the Pvu II site on the 5' side 26 of the SV40 early region promoter to produce pSVPLX.

The pCDW and pSVPLX plasmids were digested with enzymes Hind III plus Xba I from New England Biolabs and their DNAs isolated using the Geneclean product following agarose gel electrophoresis. Ligation of the CD4

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fragment into the pSVPLX vector was performed and 1 colonies were screened to yield pSVCDW in which the SV40 2 virus early region promoter is used to 3 4 expression of the complete CD4 gene product. The next step was to produce a form of the CD4 gene such that it 5 6 would be exported from the cell as an extracellular 7 product.

C. The production of a soluble form of CD4 a specially designed accomplished by the use of oligonucleotide adaptor to produce a mutant form of the 10 11 CD4 gene. This adaptor has the unique property that when inserted into the Nhe I site of the CD4 gene it produces 12 the precise premature termination of the CD4 protein 13 amino acid sequence while regenerating the Nhe I site and 14 creating a new Hpa I site. This oligonucleotide adaptor, 15 synthesized by Midland Certified Reagent Co., 16 17 produced bу annealing two phosphorylated oligonucleotides: (1) 5'CTAGCITGAGTGAGIT 3' 18 This product was then ligated into the AACTCACTCAAG. 19 site of pSVCDW. The ligation reaction was then cleaved 20 with Hpa I and then Xho I linkers were added. The linker 21 reaction was terminated by heating at 65°C for 15 minutes 22 and then subjected to digestion with Xho I restriction 23 24 endonuclease from New England Biolabs. This reaction was then subjected to agarose gel electrophoresis and the 25 fragment containing the SV40-CD4 adaptor isolated using 26 27 the Geneclean product. The retroviral vector N2 was prepared to accept the SV40-CD4-adaptor fragment by 28 digestion with Xho I and treatment with calf intestinal 29 30 phosphatase from Boehringer Mannheim, Indianapolis, 31 Indiana.

- 1 The ligation of a CD4 expression cassette was performed 2 with an insert to vector ratio of 5:1 3 transformed in DH5 competent bacteria from Bethesda 4 Research Labs. Constructs were analyzed by restriction 5 endonuclease digestion to screen for orientation and then 6 grow up in large scale. The construct where the SV40 7 virus promoter is in the same orientation as the viral 8 LTR promoters is known as SSC while the construction in 9 the reverse orientation is called SCSX.
- The SSC vector is packaged into PA 317 cell line
 as described by Miller, et al., <u>supra</u>, to provide PA 317
 cells capable of producing soluble CD4 protein. The SSC
 vector packaged PA 317 cells were used to transduce
 rabbit endothelial cells as described above. The
 transduced endothelial cells expressed soluble CD4.
- 16 D. Collagen sponges containing adsorbed HBGF-1 of the 17 type previously described were surgically implanted in 18 the abdominal cavity of a rat near the liver. 19 surgically removed seven to 20 post-implantation and digested 30 to 60 minutes at 21 with a solution of collagenase in phosphate buffered 22 saline in a concentration of lmg/ml using a tissue 23 culture incubator at 5 percent in CO2. Released cells 24 were collected by centrifugation for 10 minutes at 1000 25 RPM at 20°C. The cells were washed once with phosphate 26 buffered saline (PBS) and pelleted by centrifugation. 27 Cells were resuspended with two volumes of 30 ml of media 28 containing: M199 media (Gibco); ECGF (crude 29 extract) 7.2mg; Heparin (Upjohn) 750 units;

- and 20 percent conditioned cellular media collected as 1 supernatant from confluent dishes after 48 hours of 2 either bovine aortic or human umbilical vein endothelial 3 cells. The other media contained: 10 percent fetal calf 4 serum (Hyclone); 3000 units Penicillan G (Biofluids); and 5 3000 units streptomycin sulfate (Biofluids) and the cells 6 were plated for 16 hours on 100 mm tissue culture disk 7 coated with fibronectin (human) using lug/cm2. Plated 8 cells were washed with PBS three times and fed 15ml of 9 previously mentioned media. Media was changed every 2 10 days for the duration of the procedures.
- Selected rat endothelial cells were transduced 12 with N-7, SAX, G2N and SSC vectors by the following 13 procedures: 14
- 1. 2×10^6 microendothelial cells (monolayer 80 15 16 percent confluent)
- 2. 2 x 106 cfu/ml viral supernatant 17
- 18 Polybrene (8ug/ml)
- Combine 1, 2, 3 in 5 ml total volume for 2-3 19 hours at 37°C (5 percent CO2). 20
- Add 20ml of tissue culture media for 16 hours, 21
- at 37°C (5 percent CO2). 22
- Aspirate off media (virus containing), add 23
- 24 fresh culture media.

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1	- After 48-96 hours, add G418 (800ug/ml) and
2	culture media.
3	- Select for one to two weeks changing media
4	every two days.
5	The following are procedures for seeding a sponge
6	with the transduced endothelial cells described above

7 A. The endothelial cells are seeded directly onto a 8 HBGF-1 adsorbed, collagen coated PTFE fiber sponge, and the sponge is implanted back into the same animal used as 9 10 the source of endothelial cells. The site 11 implantation can be subcutaneous, intraperitoneal, or at 12 or near the site of the organ that normally produces the 13 new product encoded by the gene transduced into the 14 endothelial cells. The sponge implant generates its own vascularization within 5 to 10 days, as described in 15 16 earlier examples. The engineered endothelial cells are 17 maintained on the implant such that the new gene product 18 is delivered directly into the circulation after 19 secretion from the cell. The production of the gene 20 product is monitored either by direct measurement of its 21 serum levels, by the biochemical or physiological effect 22 of the agent, or both.

B. An HBGF-1 absorbed, collagen coated PTFE fiber sponge is preimplanted at the desired site, as described above, and at the time determined to be optional for that implant site for establishment of neovascularization. The transformed cells are injected directly into the

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1 already-vascularized fiber sponge. The advantage of this 2 method is that the engineered cells are more rapidly and 3 effectively, established in the implant or migrate back 4 into the parent organ (e.g., liver). The product begins 5 to enter the circulation much sooner than with method A above. Production of the new gene product is measured as 7 described in method A. This procedure can be applied to R a number of different cell types capable of being q sampled, genetically engineered in vivo, and reinserted 10 via the fiber sponge implant. Such cells 11 fibroblasts. hepatocytes, smooth muscle cells, bone 12 marrow cells and others. The products delivered to the 13 circulation can be any peptide or protein whose gene can 14 be inserted into a cell and whose product is desired to 15 be delivered.

16 EXAMPLE 7

17 Gortex shunt tubes were surgically implanted into 18 the peritoneum of rats, in such a way as to form a loop. 19 with each end contacting the aorta. The tubes contained 20 either a Gelfoam (Collagen I) sponge impregnated with 21 HBGF-1 (1 ng/ml) or a bundle of "angel hair" Gortex 22 fibers, coated with Collagen I and impregnated with 23 HBGF-1 (1 ng/ml). The tubes were left in the animals for 24 one month, then surgically extracted, grossly examined 25 for blood vessel formation, and the sponge prepared for 26 histological examination. As shown in Figure 14A, the

tube that had contained the Gelfoam sponge contained no new blood vessels, and the sponge had completely dissolved. In contrast, the angel-hair Gortex fiber bundles became significantly vascularized (Figure 14B), with higher magnification showing the capillary 6 structures (Figures 14C, D).

7 This experiment provides an example of directing neovascularization to a particular site, with a two 8 9 component device. The first component, a tube or pouch, can provide a receptacle in which implanted cells, 10 11 genetically engineered or normal, can be seeded. 12 possible that such a site may be immunologically 13 privileged, and allow cells from another individual, or 14 even another species, to survive and produce a desired product. 15

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and mixtures thereof.

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1	WHAT IS CLAIMED IS
2	1. A neovascularization device comprising
3	a biocompatible support; and
4 5	a biological response modifier for inducing neovascularization, said biological response modifier
6	being adsorbed to said biochemical support.
7 8	2. The neovascularation device of claim 1 wherein said biocompatible support is an absorbable
9 .	support.
10 11	3. The neovascularization device of claim 2 further comprising:
12	a non-absorbable support.
13	4. The neovascularization device of claim 1
14	wherein said biocompatible support is a non-absorbable
15	support.
16	5. The neovascularization device of claim 2
17	wherein said absorbable support is a member selected from
18	the group consisting of collagen, laminin, fibronecting,
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gelatin, glycosaminoglycan, glycoproteins, proteoglycans

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- 1 The neovascularization device of claim 1 2 wherein said biological response modifier is a member 3 selected from the group consisting of a hormone, a 4 hormone prototype, a hydrolase, and mixtures thereof.
- 5 7. The neovascularization device of claim 6 6 wherein said hormone is an angiogenic and neurotrophic 7 growth factor being a member selected from the group consisting of HBGF-I, HBGF-II, an HBGF-I prototype, an 8 9 HBGF-II prototype, and mixtures thereof.
- 8. The neovascularization device of claim 6 said hydrolase is heparinase, collagenase, plasmin, a plasminogen activator, thrombin, heparatinase, 13 and mixtures thereof.
 - 9. The neovascularization device of claim 1 wherein said biological response modifier angiogenic growth factor, said angiogenic growth factor being in a concentration of about 1 to about 10 nanograms per mm3 of said support.
- 19 10. The neovascularization device of claim 3 20 wherein said non-absorbable support is a member selected 21 from the group consisting of nylon, rayon, 22 polypropylene, polyethylene, PTFE, collagen I, collagen 23 IV, kerratin, and glycolipid.

	••
1 2 3 4 5	11. The neovascularization device of claim 4 wherein said non-absorbable support is a member selected from the group consisting of nylon, rayon, dacron, polypropylene, polyethylene, PTFE, collagen I, collagen IV, kerratin, and glycolipid.
6 7	12. The neovascularization device of claim 2 wherein said absorbable support is gelatin.
8	13. A neovascularization device comprising:
9	an absorbable support;
10 11	a non-absorbable support, said absorbable support being adsorbed to said non-absorbable support; and
12	a biological response modifier in sufficient

- a biological response modifier in sufficient 13 concentration for inducing in vivo site directed 14 neovascularization, said biological response modifier 15 being adsorbed to said absorbable support.
- 16 The neovascularization device of claim 13 17 wherein said absorbable support is a member selected from 18 the group consisting of collagen, laminin, fibronectins, 19 gelatin, glycosaminoglycan, glycoproteins, proteoglycans 20 and mixtures thereof.
- 21 The neovascularization device of claim 13 22 wherein said biological response modifier is a member 23 selected from the group consisting of a hormone, a 24 hormone prototype, a hydrolase, and mixtures thereof.

- 1 16. The neovascularization device of claim 15
 2 wherein said hormone is an angiogenic and neurotrophic
 3 growth factor being a member selected from the group
 4 consisting of HBGF-I, HBGF-II, an HBGF-I prototype, an
 5 HBGF-II prototype, and mixtures thereof.
- 6 17. The neovascularization device of claim 15
 7 wherein said hydrolase is heparinase, collagenase,
 9 plasmin, a plasminogen activator, thrombin, heparatinase,
 9 and mixtures thereof.
- 10 18. The neovascularization device of claim 13
 11 wherein said biological response modifier is an
 12 angiogenic growth factor, said angiogenic growth factor
 13 being in a concentration of about 1 to about 10 nanograms
 14 per mm³ of said per mm³ of both said absorbable support
 15 and non-absorbable support.
- 16 19. The neovascularization device of claim 13
 17 wherein said non-absorbable support is a member selected
 18 from the group consisting of nylon, rayon, dacron,
 19 polypropylene, polyethylene, PTFE, collagen I, collagen
 20 IV, kerratin, and glycolipid.
- 21 20. A neovascularization device comprising:
- 22 a biocompatible support; and

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and mixtures thereof.

1	a biological response modifier for inducing in
2	vivo site directed neovascularization, said biological
3	responses modifier being (i) in a concentration of about
4	1 to about 10 nangrams per mm3 of said biocompatible
5	support and (ii) a member of the group consisting of a
6	hormone, a hormone prototype, a hydrolase, and mixtures
7	thereof.
8	21. The neovascularation device of claim 20
9	wherein said biocompatible support is an absorbable
10	support.
11	22. The neovascularization device of claim 21
12	further comprising:
	•
13	a non-absorbable support.
13	a non-absorbable support.
14	23. The neovascularization device of claim 20
15	wherein said biocompatible support is a non-absorbable
16	support.
17	24. The neovascularization device of claim 21
18	wherein said absorbable support is a member selected from

the group consisting of collagen, laminin, fibronectins,

gelatin, glycosaminoglycan, glycoproteins, proteoglycans

- 1 25. The neovascularization device of claim 20
 2 wherein said hormone is an angiogenic and neurotrophic
 3 growth factor being a member selected from the group
 4 consisting of HBGF-I, HBGF-II, an HBGF-I prototype, an
 5 HBGF-II prototype, and mixtures thereof.
- 6 26. The neovascularization device of claim 20 7 wherein said hydrolase is heparinase, collagenase, plasmin, a plasminogen activator, thrombin, heparatinase, and mixtures thereof.
- 10 27. The neovascularization device of claim 22
 11 wherein said support is a member selected from the group
 12 consisting of nylon, rayon, dacron, polypropylene,
 13 polyethylene, PTFE, collagen I, collagen IV, kerratin,
 14 and glycolipid.
- 15 28. The neovascularization device of claim 23
 16 wherein said non-absorbable support is a member selected
 17 from the group consisting of nylon, rayon, dacron,
 18 polypropylene, polyethylene, PTFE, collagen I, collagen
 19 IV, kerratin, and glycolipid.
 - 20 29. A process for producing neovascularization 21 comprising:
 - 22 adsorbing a biological response modifier for 23 inducing neovascularization onto a biocompatible support;

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1	contacting a therapeutically effective amount of
2	said adsorbed biological response modifier to at least
3	one selected tissue in an organism; and
4	directing in vivo growth of neovascular cells at
5	said contacted, selected tissue for a sufficient time to
6	obtain a vascular structure.
7	30. The process for producing neovascularization
8	of claim 29 wherein said neovascular cells contain a

- 10 The process for producing neovascularization
- 11 of claim 30 wherein said genetic insert enables said 12
- neovascular cells to secrete a biological product.
- The process for producing neovascularization 13 14 of claim 31 wherein said biological product
- 15 biological response modifier.

genetic insert.

- 16 33. The process for producing neovascularization of claim 32 wherein said biological response modifier is 17
- a member selected from the group consisting of a hormone, 18
- a hormone precursor, and a hydrolase. 19
- 20 34. The process for producing neovascularization 21 of claim 29 further comprising:

cells.

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support.

seeding said vascular structure with non-vascular

3	35. The process for producing neovascularization
4	of claim 34 wherein said seeded cells secrete a desired
5	biological product.
6	36. The process for producing neovascularization
7	of claim 34 wherein said seeded cells perform a desired
8	metabolic function.
9	37. The process for producing neovascularation
10	of claim 29 wherein said biocompatible support is an
11	absorbable support.
12	38. The neovascularization device of claim 37
13	further comprising:
14	a non-absorbable support.

18 40. The neovascularization device of claim 37
19 wherein said absorbable support is a member selected from
20 the group consisting of collagen, laminin, fibronectins,
21 gelatin, glycosaminoglycan, glycoproteins, proteoglycans
22 and mixtures thereof.

39. The neovascularization device of claim 29

wherein said biocompatible support is a non-absorbable

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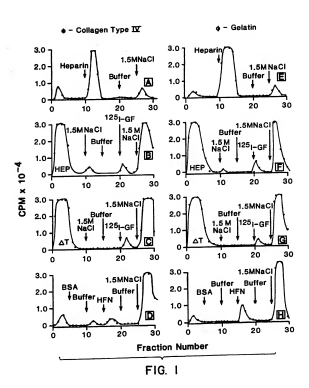
- 1 41. The neovascularization device of claim 29
 2 wherein said biological response modifier is a member
 3 selected from the group consisting of a hormone, a
 4 hormone prototype, a hydrolase, and mixtures thereof.
- 5 42. The neovascularization device of claim 41 6 wherein said hormone is an angiogenic and neurotrophic 7 growth factor being a member selected from the group 6 consisting of HBGF-I, HBGF-II, an HBGF-I prototype, an 9 HBGF-II prototype, and mixtures thereof.
- 10 43. The neovascularization device of claim 41
 11 wherein said hydrolase is heparinase, collagenase,
 12 plasmin, a plasminogen activator, thrombin, heparatinase,
 13 and mixtures thereof.
 - 44. The neovascularization device of claim 29 wherein said biological response modifier is an angiogenic growth factor, said angiogenic growth factor being in a concentration of about 1 to about 10 nanograms per mm³ of said support.
- 19 45. The neovascularization device of claim 38
 20 wherein said non-absorbable support is a member selected
 21 from the group consisting of nylon, rayon, dacron,
 22 polypropylene, polyethylene, PTFE, collagen I, collagen
 23 IV. kerratin, and divcolipid.

1	46. The neovascularization device of claim 39
2	wherein said non-absorbable support is a member selected
3	from the group consisting of nylon, rayon, dacron,
4	polypropylene, polyethylene, PTFE, collagen I, collagen
5	IV, kerratin, and glycolipid.
6 7	47. A product for promoting neovascularization, comprising:
8	a support including an extracellular matrix
9	protein and a biological response modifier.

- 10 48. The product of claim 47 wherein the support
- 11 includes cells capable of expressing a metabolite whereby
 12 the product is capable of inducing organoid
 13 neovascularization.
- 49. The product of claim 48 wherein the cells are genetically engineered to express a heterologous protein.
- 16 50. The product of claim 49 wherein the support 17 is a non-absorbable support.
- 18 51. The product of claim 50 wherein the 19 biological response modifier is absorbed to the 20 extracellular matrix protein included in the 21 non-absorbable support.

L	52.	The	product	of	claim	51	wherein	said
2	biological	response	modifier	is a	member	se	elected	from
3	the group	consisting	of a hor	mone,	a hormo	one	prototy	e, a
ı	hydrolase	and mixto	ires there	of.				

- 5 53. The product of claim 52 wherein the biological response modifier is at least one member selected from the group consisting of heparinase, collagenase, plasmin, a plasminogen activator, thrombin,
- 8 collagenase, plasmin, a plasminogen activator, thrombin 9 and heparatinase.
- 10 54. The product of claim 52 wherein the 11 biological response modifier is at least one member 12 selected from the group consisting of HBGF-I, HBGF-II, 13 and HBGF-I prototype, and an HBGF-II prototype.
- 14 55. The product of claim 51 wherein said 15 biological response modifier is an angiogenic growth 16 factor, said angiogenic growth factor being in a 17 concentration of about 1 to about 10 nanograms per mm³ of 18 said support.
- 19 56. The product of claim 51 wherein said 20 non-adsorbable support is a member selected from the 21 group consisting of nylon, rayon, daoron, polypropylene, 22 polyethylene, PTFE, and cross-linked collagen IV.
- 23 57. The product of claim 51 wherein the
 24 extracellular matrix protein is at least one member
 25 selected from the group consisting of collagen, laminin,
 26 fibronectins, gelatin, glycosaminoglycan, glycoproteins,
 27 and proteoglycans.



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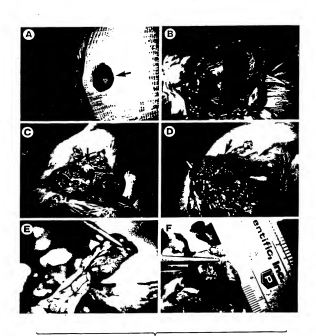


FIG. 2

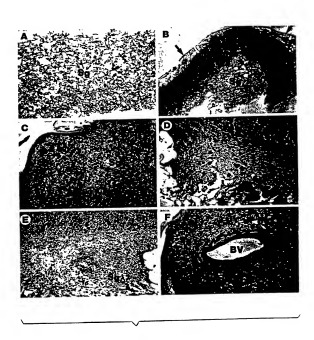
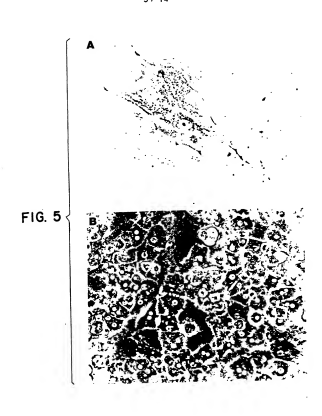


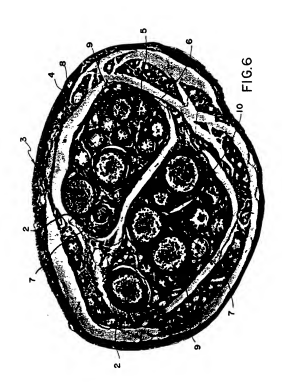
FIG. 3



FIG. 4



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16. 7



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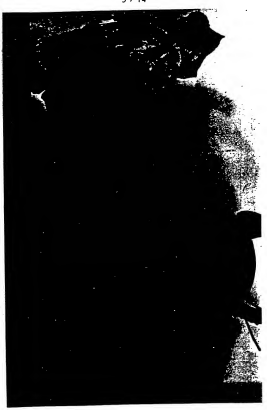
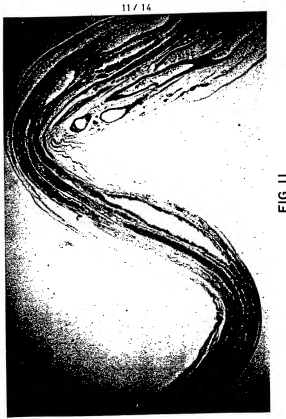


FIG. 9





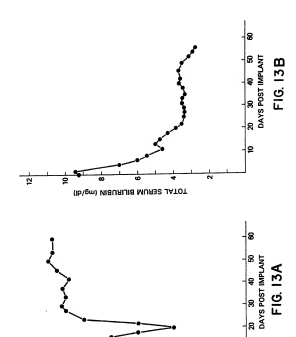


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FIG. 14

International Application No. PCT/US89/00742

I. CLASS	IFICATIO	N OF SUBJECT MATTER (II Severel classif	ication symbols apply, indicate all) *			
TDC	4) + Af	onel Patent Clessification (IPC) or to both Nation 151K 37/02,37/24,37/54;	C12N 5/00,11/02,11/	/08 177,180,240.23		
	U.S./C1.: 514/2,8;424/94.6,94.61,94.63,94.64;435/177,180,240.23					
II. FIELDS	SEARCH	Minimum Documen	tetion Searched 7			
Clessification	- Custom		Clessification Symbols			
U.S.	on System	424/94.6,94.61,94.63,	94.64;435/177,180,7	240.23		
0.0.		514/2,8,21,774,801		·		
		,_,_,_,_,				
		Documentation Searched other to	nen Minimum Documentation are Included in the Fields Searched			
_DATA	BASES	CHEMICAL ABSTRACTS	ERVICES ONLINE (FIL	e for		
		FILE BIOSIS, 1969-19	89). See Attachment	T TOE		
Sear	ch te	cms.				
III DOCI	MENTS '	ONSIDERED TO BE RELEVANT				
Category *	Citet	ion of Document, 11 with indication, where appr	opriate, of the relevant passages 12	Relevant to Cleim No. 13		
Y	US, I	A, 4,699,141 (Lamberto	n et al) 13	13-16,18,		
-	Octo	er 1987, See Entire D	ocument	19,22,27,		
		•		30-38,40,		
				42,45,		
				47-52,		
				54-57		
				1 4 20		
x	US, I	4, 4,699,141 (Lamberto	n et al) 13	1,4,29,		
	Octo	oer 1987, See Entire D	ocument	34,35,36,		
				39,47,48		
	D	7 of Dr	ontal Pathology	1,4,6,		
Y	Brit:	ish Journal of Experim	drade et al.	9-11,20,		
	volu	ne 68, Issued 1987, An	ine on	23,28,29,		
	'Qua	ntitative in Vivo Stud	go model" nages	39,41,44,		
	angi	ogenesis in a rat spon	de moner ' hades	46		
	/55-	766, Entire Document.		1		
Y	Proc	edings of the Nationa	1 Academy of	1,4,6,9,		
	Scie	nces, USA, Volume 82,	Issued November	11,20,23,		
	1985	, Buckley et al, "Sust	ained release of	28,29,39,		
	enide	ermal growth factor ac	celerates wound	41,44,45,		
	repa	ir," pages 7340-7344,	Entire Document.	46		
		, 2-3,				
				L. Carrellond Silve Con-		
* Specie	el categorie	e of cited documents: 10	"I" leter document published after to priority dete end not in conflicted to understand the princip	ict with the application but		
cor	reidered to	ning the general state of the art which is not be of particular relevence				
"E" eer	tier docume	ant but published on or efter the international	"X" document of perticular relevan	ce; the claimed invention cennot be considered to		
"" decument which may throw doubt on priority claim(a) or which is claim to action in the publication derived in another which is claim to action action are procised reason (as specified). "Of document referring to an oral disclosure, use, axhibition or document is combined with one or more other excit document."						
"O" document referring to an oral disclosure, use, exhibition or mants, such combination being obvious to a person skilled						
other meens "P" document published prior to the internetional filing data but leter than the priority date claimed "a" document member of the same patent family						
IV. CERTIFICATION Detect the Actual Completion of the International Search Date of Mailing of this International Search Report						
Dete of the Actual Congression of the International Search Dete of Mailing of this International Search Report 0 5 JUL 1989						
Internatio	nal Searchi	ng Authority	Signature of Authorized Officer			
18	A/US		Hil I Krox	2		

International	Application No.	
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FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET				
XY	Biochemical and Biophysical Research Communications Volume 147, Issued 15 September 1987 Hayek et al, "An In Vivo model for study of the angiogenic effects of basic fibroblast growth factor", pages 876-880, Entire document.	1-7, 12-16, 29, 37-42 9-11, 18-25, 27-28, 44-46		
V. 🗌 08	SERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE!			
Tile international search report has not been established in respect of certain claims under Article 17(0) (o) for the following reasons: Claim numbers				
Claim numbers				
VI. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING?				
This toternational Searching Authority found multiple invantions in this international application as tollows:				
1. A	all required additional search less were timely paid by the applicant, this international search raport of	overs all searchable claims		
1 . m	of the international application. 2. A only once of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of this international application for which fees were paid, specifically failims:			
3. No	required additional search fees were timely paid by the spolicant. Consequently, this international se invention first mentioned in the claims; it is covered by claim numbers:	arch report is restricted to		
Remark	s all searchable claims could be searched without effort justifying an additional fee, the International is nis payment of any additional lee. on Protest	Searching Authority did not		
	ne additional search fees were accompanied by applicant's protest.			
<u></u>	o protest accompanied the payment of additional search fees.			

III. DOCUMENTS CONSIDERED TO SE RELEVANT (CONTINUED FROM THE SECOND SHEET)		
Category *	Citation of Document, 16 with Indication, where eppropriate, of the relevant passages 17	Relevant to Claim No 1
Y	WO, A 8701728 (Biotechnology Research Partners, LTD) 26 March 1987, pages 1-64, See pages 1-3, 12-16.	1,4,6, 7,9,11, 20,23, 25,28, 29,39, 44,46
х,Р	Journal of Cell Biology, Volume 107 (5 part 3), Issued 09 December 1988, Maciag et al, "Heparin-Binding Growth Factor-I (HBGF-I) Bound to Gelatin Induces Site-Specfic Neovessel Formation In Vivo", Abstract No. 2697, page 479a.	1-57
Υ	Surgical Science Series, Volume 2, Issued 1984, Jackson et al, "Effect of Angiogenic Factors on the Vascularization of IVALON Sponge Implants", pages 190-201, See entire document.	1,4,6, 7,9, 11,20, 23,25, 28,29 39,44,
Υ	Biochemical and Biophysical Research Communications, Volume 142, Issued 30 January 1987, Baird et al, "Fibroblast Browth Factors are Present in the Extracellular Matrix Produced by Endothebal Cells In Vitro. Implications for a Role of heporinase-like Enzymes in the Neovascular Response, pages 428-435, especially page 433.	8,17, 26,43, 53
;	Cancer Research, Volume 43, Issued June 1983, Vlodavsky et al, "Lymphona Cell-Mediated Degradation of sulfated Proteoglycans in the Subendothelial Extracellular Matrix: Relationship to Tumor Cell Metastiasis", pages 2704-2711, especially page 2710.	26,43,
Y	Chemical Abstracts, Volume 109, Issued 10 October 1988, Rifkin, "Endothelial Cell proteases and Cellular invasion", Abstract No.:123103	8,17, 26,43 53

Attachment to Form PCT/ISA/210 Part II. FIELDS SEARCHED SEARCH TERMS:

Angiogenesis
Neovascularization
IVALON
Neovessel
formation
sponge
support
biocompatible
transplant
heparinase
heparitinase
collagenase
plasminogen
activator
plasmin
hydrolase